

STIC Search Report Biotech-Chem Library

STIC Database Tracking Number: 131201

TO: Howard Owens Location: 5d34 / 5c18 Monday, August 30, 2004

Art Unit: 1623 Phone: 272-0658

Serial Number: 10 / 602976

From: Jan Delaval

Location: Biotech-Chem Library

Rem 1A51

Phone: 272-2504

jan.delaval@uspto.gov

Search Notes		And the second s	
	-		
			e:
			<u> </u>



=> fil req

FILE 'REGISTRY' ENTERED AT 14:41:34 ON 30 AUG 2004
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
COPYRIGHT (C) 2004 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 29 AUG 2004 HIGHEST RN 735258-95-4 DICTIONARY FILE UPDATES: 29 AUG 2004 HIGHEST RN 735258-95-4

TSCA INFORMATION NOW CURRENT THROUGH MAY 21, 2004

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

=> d sta que 123

L20 STF

VAR G1=O/S/SO2/CH2
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT IS PCY AT 6
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS M3 N AT 6

GRAPH ATTRIBUTES:

RSPEC 1

NUMBER OF NODES IS 8

STEREO ATTRIBUTES: NONE

L21 12715 SEA FILE=REGISTRY ABB=ON PLU=ON (N2C3-NCNC3 OR NCNC2-NC5 OR N3C2-NC5 OR N2CNC-NC5)/ES AND (OC4 OR SC4 OR C5)/ES

L23 2684 SEA FILE=REGISTRY SUB=L21 SSS FUL L20

100.0% PROCESSED 2966 ITERATIONS

2684 ANSWERS

SEARCH TIME: 00.00.01

=> d his

L1

(FILE 'HOME' ENTERED AT 14:02:39 ON 30 AUG 2004) SET COST OFF

FILE 'HCAPLUS' ENTERED AT 14:02:47 ON 30 AUG 2004

1 S (US20040101535 OR US20030050229)/PN OR US2000-206585#/AP,PRN
E SOMMADOSSI J/AU

```
L2
            210 S E4, E5
                E LACOLLA P/AU
L3
              5 S E4,E5
                E LA COLLA P/AU
            194 S E3-E7
                E COLLA /AU
                E NOVIRIO/PA, CS
L5
              14 S E3-E17
                SEL RN L1
     FILE 'REGISTRY' ENTERED AT 14:04:34 ON 30 AUG 2004
L6
             18 S E1-E18
     FILE 'HCAPLUS' ENTERED AT 14:06:43 ON 30 AUG 2004
            402 S L2-L5 NOT L1
L7
     FILE 'REGISTRY' ENTERED AT 14:07:30 ON 30 AUG 2004
     FILE 'HCAPLUS' ENTERED AT 14:07:30 ON 30 AUG 2004
                SET SMARTSELECT ON
1.8
            SEL L7 1- RN :
                              5164 TERMS
                SET SMARTSELECT OFF
     FILE 'REGISTRY' ENTERED AT 14:07:45 ON 30 AUG 2004
           5164 S L8
L9
           1007 S L9 AND OC4/ES
L10
            646 S C2N3-C5N/EA AND OC4/ES
L11
L12
           3462 S C3N2-C5N/EA AND OC4/ES
           2889 S NCNC2-NC5/ES AND OC4/ES
L13
            573 S L12 NOT L13
L14
L15
            497 S N2CNC-NC5/ES AND L11
            149 S N3C2-NC5/ES AND L11
L16
L17
            646 S L15, L16
           3655 S N2C3-NCNC3/ES AND OC4/ES
L18
             76 S L9 AND L13, L17, L18
L19
L20
                STR
          12715 S (N2C3-NCNC3 OR NCNC2-NC5 OR N3C2-NC5 OR N2CNC-NC5)/ES AND (OC
L21
L22
             50 S L20 SAM SUB=L21
           2684 S L20 FUL SUB=L21
L23
                SAV TEMP L23 OWENS602/A
             55 S L9 AND L23
L24
L25
             21 S L19 NOT L24
L26
           2629 S L23 NOT L24
     FILE 'HCAPLUS' ENTERED AT 14:24:08 ON 30 AUG 2004
L27
            287 S L24
L28
           1420 S L26
L29
           1434 S L27, L28 AND (PD<=20000523 OR PRD<=20000523 OR AD<=20000523)
                E HEPATITIS C/CT
L30
           6755 S E5-E15
                E E5+ALL
L31
           8412 S E8, E9, E6+NT
                E HEPATITIS C/CT
                E E3+ALL
           4192 S E2
L32
          11890 S HEPATITIS C
L33
              1 S L29 AND L30-L33
L34
                E HEPATITIS/CT
          16169 S E3-E28
L35
           1425 S E34+OLD, NT, PFT, RT
L36
           9616 S E52+OLD, NT, PFT, RT
L37
            754 S E76+OLD, NT, PFT, RT
L38
```

485 S E78+OLD, NT, PFT, RT

L39

```
L40
             15 S E79+OLD, NT, PFT, RT
L41
            561 S E81+OLD, NT, PFT, RT
L42
            512 S E85+OLD, NT, PFT, RT
L43
           7978 S E90+OLD, NT, PFT, RT
                E E3+ALL
                E HEPATITIS VIRUS/CT
                E E3+ALL
L44
           7978 S E4,E3
L45
          43705 S HEPATITIS
L46
              6 S L29 AND L35-L45
L47
              3 S L1-L5 AND L27, L28
L48
              9 S L34, L46, L47
L49
              6 S L29 AND ?HEPATITIS?
L50
              9 S L48, L49
     FILE 'REGISTRY' ENTERED AT 14:30:02 ON 30 AUG 2004
              6 S (RIBAVIRIN OR PROTEASE OR POLYMERASE OR HELICASE)/CN
L51
     FILE 'HCAPLUS' ENTERED AT 14:30:39 ON 30 AUG 2004
L52
             37 S L51 AND L29
             64 S (RIBAVIRIN OR PROTEASE OR PROTEINASE OR POLYMERASE OR HELICAS
L53
              6 S INTERFERON AND L29
L54
     FILE 'REGISTRY' ENTERED AT 14:31:29 ON 30 AUG 2004
              1 S THIAZOLIDINE/CN
L55
     FILE 'HCAPLUS' ENTERED AT 14:31:32 ON 30 AUG 2004
L56
              0 S L55 AND L29
              0 S THIAZOLIDIN? AND L29
L57
L58
             80 S L52-L54
L59
             16 S L58 AND (MIX? OR COMBIN? OR SYNERG? OR COMPOSITION OR COTHERA
                SEL DN AN 13 L59
L60
              1 S L59 AND E1-E3
L61
             64 S L58 NOT L59
                SEL DN AN 14 18 25
L62
              3 S L61 AND E4-E12
L63
              1 S L62 AND HEPATITIS
L64
             10 S L50, L60, L63
             10 S L64 AND L1-L5, L27-L50, L52-L54, L56-L64
L65
                SEL HIT RN
     FILE 'REGISTRY' ENTERED AT 14:40:10 ON 30 AUG 2004
L66
             77 S E13-E89
L67
              1 S L66 AND L51
L68
             76 S L66 AND L23
     FILE 'REGISTRY' ENTERED AT 14:41:34 ON 30 AUG 2004
=> d ide can 167
     ANSWER 1 OF 1 REGISTRY COPYRIGHT 2004 ACS on STN
L67
     36791-04-5 REGISTRY
RN
     1H-1,2,4-Triazole-3-carboxamide, 1-β-D-ribofuranosyl- (9CI)
                                                                    (CA
CN
     INDEX NAME)
OTHER NAMES:
     1-\beta-D-Ribofuranosyl-1,2,4-triazol-3-carboxyamide
CN
     1-β-D-Ribofuranosyl-1,2,4-triazole-3-carboxamide
CN
CN
     ICN 1229
     NSC 163039
CN
CN
     Ravanex
CN
     Rebetol
CN
     Ribamide
```

CN

Ribamidil

CN Ribavarin

CN Ribavirin

CN Tribavirin

CN Vilona

CN Viramid

CN Virazole

FS STEREOSEARCH

DR 66510-90-5, 437710-49-1

MF C8 H12 N4 O5

CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN*, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, CSNB, DDFU, DIOGENES, DRUGU, EMBASE, HSDB*, IFICDB, IFIPAT, IFIUDB, IMSCOSEARCH, IMSDRUGNEWS, IMSPATENTS, IPA, MEDLINE, MRCK*, MSDS-OHS, NAPRALERT, NIOSHTIC, PHAR, PIRA, PROMT, PROUSDDR, PS, RTECS*, SYNTHLINE, TOXCENTER, USAN, USPAT2, USPATFULL, VETU

(*File contains numerically searchable property data)

Other Sources: WHO

DT.CA CAplus document type: Book; Conference; Dissertation; Journal; Patent; Report

RL.P Roles from patents: ANST (Analytical study); BIOL (Biological study); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)

RLD.P Roles for non-specific derivatives from patents: BIOL (Biological study); PREP (Preparation); USES (Uses)

RL.NP Roles from non-patents: ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative); MSC (Miscellaneous); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)

RLD.NP Roles for non-specific derivatives from non-patents: BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1977 REFERENCES IN FILE CA (1907 TO DATE)
71 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
1985 REFERENCES IN FILE CAPLUS (1907 TO DATE)

REFERENCE 1: 141:162353

REFERENCE 2: 141:155723

REFERENCE 3: 141:140707

REFERENCE 4: 141:138913

REFERENCE 5: 141:133747

REFERENCE 6: 141:133621

REFERENCE 7: 141:128820

REFERENCE 8: 141:128600

REFERENCE 9: 141:122155

REFERENCE 10: 141:122153

=> fil hcaplus

FILE 'HCAPLUS' ENTERED AT 14:41:56 ON 30 AUG 2004
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications. The CA Lexicon is the copyrighted intellectual property of the the American Chemical Society and is provided to assist you in searching databases on STN. Any dissemination, distribution, copying, or storing of this information, without the prior written consent of CAS, is strictly prohibited.

FILE COVERS 1907 - 30 Aug 2004 VOL 141 ISS 10 FILE LAST UPDATED: 29 Aug 2004 (20040829/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d 165 all hitstr tot

L65 ANSWER 1 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 2000:41758 HCAPLUS

DN 132:194596

ED Entered STN: 18 Jan 2000

TI Synthesis and biological activity of 2'-fluoro-Darabinofuranosylpyrazolo[3,4-d]pyrimidine nucleosides

- AU Shortnacy-Fowler, Anita T.; Tiwari, Kamal N.; Montgomery, John A.; Buckheit, Robert W., Jr.; Secrist, John A., III; Seela, Frank
- CS Southern Research Institute, Birmingham, AL, 35255-5305, USA
- SO Helvetica Chimica Acta (1999), 82(12), 2240-2245 CODEN: HCACAV; ISSN: 0018-019X
- PB Verlag Helvetica Chimica Acta
- DT Journal
- LA English
- CC 33-9 (Carbohydrates)
 Section cross-reference(s): 1
- Coupling of 2-fluoro-3,5-di-O-benzoyl- α -D-arabinofuranosyl bromide with 4-methoxypyrazolo[3,4-d]pyrimidine gave an α -D/ β -D mixture of N1-and N2-coupled products. All the anomers were separated and deblocked to yield the corresponding nucleosides. The β -D-anomer was converted to the 4-amino derivative, which was deaminated by adenosine deaminase to give the 4-oxo compound 1-(2-Deoxy-2-fluoro- β -D-arabinofuranosyl)-4-methoxy-1H-pyrazolo[3,4-d]pyrimidine showed significantly activity against human cytomegalovirus and hepatitis B virus; its 4-amino analog showed activity against human herpes virus 8. All the compds. were non-cytotoxic in several human tumor-cell lines in culture.
- ST pyrazolo pyrimidine nucleoside prepn virucide; fluoro arabinofuranosyl pyrazolopyrimidine nucleoside prepn antiviral antitumor

```
IT
     Antitumor agents
     Antiviral agents
     Cytotoxicity
        (synthesis and biol. activity of fluoro-D-arabinofuranosyl
        pyrazolopyrimidine nucleosides)
     Nucleosides, preparation
IT
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); BIOL (Biological
     study); PREP (Preparation)
        (synthesis and biol. activity of fluoro-D-arabinofuranosyl
        pyrazolopyrimidine nucleosides)
IT
     259738-10-8P 259738-11-9P 259738-12-0P
     259738-13-1P 259738-14-2P
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); BIOL (Biological
     study); PREP (Preparation)
        (synthesis and biol. activity of fluoro-D-arabinofuranosyl
        pyrazolopyrimidine nucleosides)
IT
     5399-93-9
                  97614-44-3
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (synthesis and biol. activity of fluoro-D-arabinofuranosyl
        pyrazolopyrimidine nucleosides)
     259738-06-2P 259738-07-3P 259738-08-4P
IT
     259738-09-5P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (synthesis and biol. activity of fluoro-D-arabinofuranosyl
        pyrazolopyrimidine nucleosides)
              THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
        20
RE
(1) Brockman, R; Biochem Pharmacol 1977, V26, P2193 HCAPLUS
(2) Carson, D; Proc Natl Acad Sci U S A 1980, V77, P6065
(3) Jungmann, O; Tetrahedron Lett 1996, V37, P8355 HCAPLUS
(4) Kazimierczuk, Z; J Am Chem Soc 1984, V106, P6379 HCAPLUS
(5) Korba, B; Antiviral Res 1992, V19, P55 HCAPLUS(6) Montgomery, J; Antimetabolites in `Cancer Chemotherapeutic Agents,' 1995,
    P47
(7) Montgomery, J; J Med Chem 1992, V35, P397 HCAPLUS
(8) Parker, W; Mol Pharmacol 1999, V55, P515 HCAPLUS (9) Robins, R; J Am Chem Soc 1956, V78, P784 HCAPLUS
(10) Secrist, J; J Med Chem 1998, V41, P3865 HCAPLUS
(11) Seela, F; Helv Chim Acta 1985, V68, P563 HCAPLUS
(12) Seela, F; Helv Chim Acta 1993, V76, P1450 HCAPLUS
(13) Seela, F; J Org Chem 1983, V48, P3119
(14) Seela, F; Nucleosides Nucleotides 1991, V10, P713 HCAPLUS
(15) Shaw, R; Cancer 1960, V13, P482 HCAPLUS
(16) Shoemaker, R; submitted
(17) Skipper, H; Cancer Res 1957, V17, P579 HCAPLUS
(18) Tann, C; J Org Chem 1985, V50, P3644 HCAPLUS
(19) Tatarowicz, W; J Virol Methods 1991, V35, P207 HCAPLUS
(20) Wright, S; Blood Rev 1994, V8, P125 MEDLINE
     259738-10-8P 259738-11-9P 259738-12-0P
     259738-13-1P 259738-14-2P
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); BIOL (Biological
     study); PREP (Preparation)
         (synthesis and biol. activity of fluoro-D-arabinofuranosyl
        pyrazolopyrimidine nucleosides)
RN
     259738-10-8 HCAPLUS
     1H-Pyrazolo[3,4-d]pyrimidine, 1-(2-deoxy-2-fluoro-β-D-
CN
     arabinofuranosyl)-4-methoxy- (9CI) (CA INDEX NAME)
```

RN 259738-11-9 HCAPLUS

CN 1H-Pyrazolo[3,4-d]pyrimidine, 1-(2-deoxy-2-fluoro- α -D-arabinofuranosyl)-4-methoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 259738-12-0 HCAPLUS

CN 2H-Pyrazolo[3,4-d]pyrimidine, 2-(2-deoxy-2-fluoro-β-D-arabinofuranosyl)-4-methoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 259738-13-1 HCAPLUS

CN 2H-Pyrazolo[3,4-d]pyrimidine, 2-(2-deoxy-2-fluoro-α-D-arabinofuranosyl)-4-methoxy- (9CI) (CA INDEX NAME)

RN 259738-14-2 HCAPLUS

CN 1H-Pyrazolo[3,4-d]pyrimidin-4-amine, 1-(2-deoxy-2-fluoro-β-D-arabinofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 259738-06-2P 259738-07-3P 259738-08-4P 259738-09-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(synthesis and biol. activity of fluoro-D-arabinofuranosyl pyrazolopyrimidine nucleosides)

RN 259738-06-2 HCAPLUS

CN 1H-Pyrazolo[3,4-d]pyrimidine, 1-(3,5-di-O-benzoyl-2-deoxy-2-fluoro-β-D-arabinofuranosyl)-4-methoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 259738-07-3 HCAPLUS

CN 1H-Pyrazolo[3,4-d]pyrimidine, 1-(3,5-di-O-benzoyl-2-deoxy-2-fluoro- α -D-arabinofuranosyl)-4-methoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 259738-08-4 HCAPLUS

CN 2H-Pyrazolo[3,4-d]pyrimidine, 2-(3,5-di-O-benzoyl-2-deoxy-2-fluoro-β-D-arabinofuranosyl)-4-methoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 259738-09-5 HCAPLUS

CN 2H-Pyrazolo[3,4-d]pyrimidine, 2-(3,5-di-O-benzoyl-2-deoxy-2-fluoro- α -D-arabinofuranosyl)-4-methoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L65 ANSWER 2 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN AN 1995:49285 HCAPLUS

```
DN 122:240293
```

ED Entered STN: 08 Nov 1994

TI 8-Aza-1-deazapurine nucleosides as antiviral agents

AU Franchetti, P.; Messini, L.; Cappellacci, L.; Abu Sheikha, G.; Grifantini, M.; Guarracino, P.; De Montis, A.; Loi, A. G.; Marongiu, M. E.; La Colla, P.

CS Dip. Sci. Chim., Univ. Camerino, Camerino, 62032, Italy

SO Nucleosides & Nucleotides (1994), 13(8), 1739-55 CODEN: NUNUD5; ISSN: 0732-8311

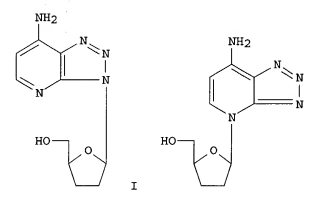
DT Journal

LA English

CC 33-9 (Carbohydrates)

Section cross-reference(s): 1

GΙ



AB Azadeazapurine nucleosides, e.g. I and II, were prepared via glycosidation of nucleobases. These dideoxy nucleosides and a series of previously synthesized 8-aza-1-deazapurine nucleosides were tested for activity against several DNA and RNA viruses, HIV-1 included. The $\alpha-$ and $\beta-$ anomers of 2',3'-dideoxy-8-aza-1-deazaadenosine were found active as inhibitors of adenosine deaminase.

TT

ST azadeazapurine nucleoside prepn virucide; dideoxyazadeazaadenosine inhibitor adenosine deaminase

IT Virucides and Virustats

(preparation and antiviral activity of azadeazapurine nucleosides)

IT Nucleosides, preparation

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and antiviral activity of azadeazapurine nucleosides)

IT 14432-09-8 34625-29-1 34641-28-6

34664-98-7 142591-84-2 142591-89-7

142591-90-0 142591-95-5 142592-00-5

162181-03-5

IT

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(preparation and antiviral activity of azadeazapurine nucleosides)

162181-09-1P 162181-11-5P 162299-76-5P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and antiviral activity of azadeazapurine nucleosides)

IT 9026-93-1, Adenosine deaminase

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(preparation and antiviral activity of azadeazapurine nucleosides)

IT 34550-49-7 127306-45-0 162299-72-1 162195-90-6 RL: RCT (Reactant); RACT (Reactant or reagent) (preparation and antiviral activity of azadeazapurine nucleosides) IT 162181-04-6P 162181-05-7P 162181-06-8P 162181-07-9P 162181-10-4P 162299-73-2P 162299-74-3P 162299-77-6P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and antiviral activity of azadeazapurine nucleosides) IT 162181-08-0P 162299-75-4P 162299-78-7P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and antiviral activity of azadeazapurine nucleosides) IT 14432-09-8 34625-29-1 34641-28-6 34664-98-7 142591-84-2 142591-89-7 142591-90-0 142591-95-5 142592-00-5 162181-03-5 RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study) (preparation and antiviral activity of azadeazapurine nucleosides) 14432-09-8 HCAPLUS RNCN 3H-Imidazo[4,5-b]pyridin-7-amine, $3-\beta$ -D-ribofuranosyl- (9CI) INDEX NAME)

Absolute stereochemistry.

RN 34625-29-1 HCAPLUS CN 2H-1,2,3-Triazolo[4,5-b]pyridine, 7-chloro-2- β -D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Absolute stereochemistry.

RN 142591-84-2 HCAPLUS
CN 3H-1,2,3-Triazolo[4,5-b]pyridin-7-amine, 3-(2-deoxy-β-D-erythro-pentofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 142591-89-7 HCAPLUS
CN 3H-1,2,3-Triazolo[4,5-b]pyridine, 7-chloro-3-(2-deoxy-β-D-erythro-pentofuranosyl)- (9CI) (CA INDEX NAME)

RN 142591-90-0 HCAPLUS

CN 3H-1,2,3-Triazolo[4,5-b]pyridine, 7-chloro-3-(2-deoxy- α -D-erythropentofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 142591-95-5 HCAPLUS

CN 2H-1,2,3-Triazolo[4,5-b]pyridin-7-amine, 2-(2-deoxy-β-D-erythropentofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 142592-00-5 HCAPLUS

CN 2H-1,2,3-Triazolo[4,5-b] pyridin-7-amine, $2-\beta-D-ribofuranosyl-$ (9CI) (CA INDEX NAME)

RN 162181-03-5 HCAPLUS CN 1H-1,2,3-Triazolo[4,5-b]pyridine, 7-chloro-1- β -D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 162181-09-1P 162181-11-5P 162299-76-5P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and antiviral activity of azadeazapurine nucleosides)

RN 162181-09-1 HCAPLUS

CN 2-Furanmethanol, 5-(7-amino-3H-1,2,3-triazolo[4,5-b]pyridin-3-yl)tetrahydro-, (2S-trans)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 162181-11-5 HCAPLUS

CN 2-Furanmethanol, 5-(7-amino-4H-1,2,3-triazolo[4,5-b]pyridin-4-yl)tetrahydro-, (2S-cis)- (9CI) (CA INDEX NAME)

RN 162299-76-5 HCAPLUS
CN 2-Furanmethanol, 5-(7-amino-3H-1,2,3-triazolo[4,5-b]pyridin-3yl)tetrahydro-, (2S-cis)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Absolute stereochemistry.

RN 162181-05-7 HCAPLUS
CN 2H-1,2,3-Triazolo[4,5-b]pyridine, 7-chloro-2-[5-[[[(1,1-dimethylethyl)diphenylsilyl]oxy]methyl]tetrahydro-2-furanyl]-, (2R-cis)-(9CI) (CA INDEX NAME)

RN 162181-06-8 HCAPLUS

CN 1H-1,2,3-Triazolo[4,5-b]pyridine, 7-chloro-1-[5-[[[(1,1-dimethylethyl)diphenylsilyl]oxy]methyl]tetrahydro-2-furanyl]-, (2S-trans)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 162181-07-9 HCAPLUS

CN 2-Furanmethanol, 5-(7-chloro-3H-1,2,3-triazolo[4,5-b]pyridin-3-yl)tetrahydro-, (2S-trans)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 162181-10-4 HCAPLUS

CN 4H-1,2,3-Triazolo[4,5-b]pyridin-7-amine, 4-[5-[[[(1,1-dimethylethyl)diphenylsilyl]oxy]methyl]tetrahydro-2-furanyl]-, (2R-cis)-(9CI) (CA INDEX NAME)

RN 162299-73-2 HCAPLUS
CN 3H-1,2,3-Triazolo[4,5-b]pyridine, 7-chloro-3-[5-[[[(1,1-dimethylethyl)diphenylsilyl]oxy]methyl]tetrahydro-2-furanyl]-, (2R-cis)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 162299-74-3 HCAPLUS
CN 2-Furanmethanol, 5-(7-chloro-3H-1,2,3-triazolo[4,5-b]pyridin-3-yl)tetrahydro-, (2S-cis)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 162299-77-6 HCAPLUS
CN 4H-1,2,3-Triazolo[4,5-b]pyridin-7-amine, 4-[5-[[[(1,1-dimethylethyl)diphenylsilyl]oxy]methyl]tetrahydro-2-furanyl]-, (2S-trans)(9CI) (CA INDEX NAME)

IT 162181-08-0P 162299-75-4P 162299-78-7P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and antiviral activity of azadeazapurine nucleosides)

RN 162181-08-0 HCAPLUS

CN 2-Furanmethanol, 5-(7-chloro-2H-1,2,3-triazolo[4,5-b]pyridin-2-yl)tetrahydro-, (2S-cis)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 162299-75-4 HCAPLUS

CN 2-Furanmethanol, 5-(7-chloro-2H-1,2,3-triazolo[4,5-b]pyridin-2-yl)tetrahydro-, (2S-trans)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 162299-78-7 HCAPLUS

CN 2-Furanmethanol, 5-(7-amino-4H-1,2,3-triazolo[4,5-b]pyridin-4-yl)tetrahydro-, (2S-trans)- (9CI) (CA INDEX NAME)

```
L65 ANSWER 3 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN
AN
     1994:473046 HCAPLUS
DN
     121:73046
ED
     Entered STN: 20 Aug 1994
     8-Aza derivatives of 3-deazapurine nucleosides. Synthesis and in vitro
ΤI
     evaluation of antiviral and antitumor activity
ΑU
     Franchetti, P.; Messini, L.; Cappellacci, L.; Grifantini, M.; Nocentini,
     G.; Guarracino, P.; Marongiu, M. E.; La Colla, P.
     Dip. Sci. Chim., Univ. Camerino, Camerino, 62032, Italy
CS
SO
     Antiviral Chemistry & Chemotherapy (1993), 4(6), 341-52
     CODEN: ACCHEH; ISSN: 0956-3202
DT
     Journal
LA
     English
CC
     1-3 (Pharmacology)
     Section cross-reference(s): 33
     The syntheses of 4-amino-1-(\beta-D-ribofuranosyl)-1H-1,2,3-triazolo[4,5-
AB
     c]pyridine (8-aza-3-deazaadenosine), 4-amino-1-(2-deoxy-\beta-D-erythro-
     pentofuranosyl)-1H-1,2,3-triazolo[4,5-c]pyridine (2'-deoxy-8-aza-3-
     deazaadenosine), and their N8 and N7 glycosylated analogs and
     4-amino-1-(2,3-dideoxy-β-D-erythro-pentofuranosyl)-1H-1,2,3-triazolo
     [4,5-c]pyridine (2',3'-dideoxy-8-aza-3-deazaadenosine) were carried out by
     glycosylation of the 4-chloro-3H-1,2,3-triazolo[4,5-c]pyridine anion. The
     anomeric configuration as well as the position of glycosylation were determined
     by 1H-, 13C-NMR, UV and N.O.E. difference spectroscopy.
     2'-Deoxy-8-aza-3-deazaadenosine and its parent compound 2'-deoxy-3-
     deazaadenosine were found active against ASFV and VSV. The
     4-chloro-2-(\beta-D-ribofuranosyl)-2H-1,2,3-triazolo[4,5-c] pyridine was
     active against Coxsackie B1, whereas none of the 8-aza-3-deaza purine
     nucleosides, compound included, was active against HIV-1. The 6-chloro
     derivs. of 8-aza-3-deazapurine ribo- and 2'-deoxyribonucleosides and
     showed some activity against LoVo human colon adenocarcinoma.
ST
     deazapurine nucleoside prepn antiviral antitumor structure; antiviral
     deazapurine nucleoside prepn structure activity; antitumor deazapurine
     nucleoside prepn structure activity
     Neoplasm inhibitors
IT
     Virucides and Virustats
        (deazapurine nucleosides, preparation and structure-activity relations of)
     Molecular structure-biological activity relationship
IT
        (neoplasm-inhibiting, of deazapurine nucleosides)
IT
     Molecular structure-biological activity relationship
        (virucidal, of deazapurine nucleosides)
IT
     36258-82-9
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (qlycosylation of)
                                            135092-93-2
IT
     4330-21-6
               16205-59-7
                              134965-83-6
     RL: BIOL (Biological study)
```

```
(glycosylation of chlorotriazolopyridine with)
IT
     154707-55-8
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (glycosylation of, with deoxydi(toluoyl)pentofuranosyl chloride)
     57680-38-3P 57680-40-7P 57680-44-1P
TТ
     154707-46-7P 154707-47-8P
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation and antitumor and antiviral activity and reaction with liquid
        ammonia)
     110483-88-0P 154707-38-7P, 2'-Deoxy-8-aza-3-deazaadenosine
IT
     154707-39-8P 154707-40-1P 154707-41-2P
     154707-48-9P
    RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); BIOL (Biological
    study); PREP (Preparation)
        (preparation and antitumor and antiviral activity of)
IT
     57680-35-0P 57680-36-1P 57680-37-2P
     154707-42-3P 154707-43-4P 154707~44-5P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and deprotection of)
TT
    154707-49-0P 154707-50-3P 154707-51-4P
     154707-52-5P 154707-53-6P 154707-54-7P
     154801-75-9P 154801-76-0P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of)
IT
    154707-45-6
     RL: BIOL (Biological study)
        (preparation reaction with liquid ammonia)
     57680-38-3P 57680-40-7P 57680-44-1P
IT
    154707-46-7P 154707-47-8P
    RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation and antitumor and antiviral activity and reaction with liquid
        ammonia)
RN
    57680-38-3 HCAPLUS
CN
    2H-1,2,3-Triazolo[4,5-c]pyridine, 4-chloro-2-β-D-ribofuranosyl- (9CI)
       (CA INDEX NAME)
```

Absolute stereochemistry.

RN 57680-44-1 HCAPLUS CN 1H-1,2,3-Triazolo[4,5-c]pyridine, 4-chloro-1-β-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 154707-46-7 HCAPLUS
CN 3H-1,2,3-Triazolo[4,5-c]pyridine, 4-chloro-3-(2-deoxy-β-D-erythro-pentofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 154707-47-8 HCAPLUS
CN 1H-1,2,3-Triazolo[4,5-c]pyridine, 4-chloro-1-(2-deoxy-β-D-erythro-pentofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 154707-39-8 HCAPLUS
CN 2-Furanmethanol, 5-(4-amino-1H-1,2,3-triazolo[4,5-c]pyridin-1-yl)tetrahydro-, (2S-cis)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 154707-40-1 HCAPLUS CN 2H-1,2,3-Triazolo[4,5-c]pyridin-4-amine, 2-β-D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 154707-48-9 HCAPLUS
CN 2H-1,2,3-Triazolo[4,5-c]pyridin-4-amine, 2-(2-deoxy-β-D-erythro-pentofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 57680-36-1 HCAPLUS

CN 2H-1,2,3-Triazolo[4,5-c]pyridine, 4-chloro-2-(2,3,5-tri-O-benzoyl-β-D-ribofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 57680-37-2 HCAPLUS

CN 3H-1,2,3-Triazolo[4,5-c]pyridine, 4-chloro-3-(2,3,5-tri-O-benzoyl- β -D-ribofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 154707-42-3 HCAPLUS

CN 2H-1,2,3-Triazolo[4,5-c]pyridine, 4-chloro-2-[2-deoxy-3,5-bis-O-(4-methylbenzoyl)-β-D-erythro-pentofuranosyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & \\ & & & \\ & & \\ & & \\ & & & \\ & & \\ & & & \\ & & \\ & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 154707-43-4 HCAPLUS

CN 3H-1,2,3-Triazolo[4,5-c]pyridine, 4-chloro-3-[2-deoxy-3,5-bis-O-(4-methylbenzoyl)-β-D-erythro-pentofuranosyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 154707-44-5 HCAPLUS

CN 1H-1,2,3-Triazolo[4,5-c]pyridine, 4-chloro-1-[2-deoxy-3,5-bis-O-(4-methylbenzoyl)-β-D-erythro-pentofuranosyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 154707-50-3 HCAPLUS
CN 1H-1,2,3-Triazolo[4,5-c]pyridine, 4-chloro-1-[5-[[[(1,1-dimethylethyl)diphenylsilyl]oxy]methyl]tetrahydro-2-furanyl]-, (2S-trans)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 154707-51-4 HCAPLUS
CN 2-Furanmethanol, 5-(4-chloro-1H-1,2,3-triazolo[4,5-c]pyridin-1yl)tetrahydro-, (2S-trans)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 154707-52-5 HCAPLUS
CN 2H-1,2,3-Triazolo[4,5-c]pyridine, 4-methoxy-2-β-D-ribofuranosyl(9CI) (CA INDEX NAME)

RN 154707-53-6 HCAPLUS
CN 1H-1,2,3-Triazolo[4,5-c]pyridine, 4-methoxy-1-β-D-ribofuranosyl(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 154707-54-7 HCAPLUS
CN 1H-1,2,3-Triazolo[4,5-c]pyridine, 1-(2-deoxy-β-D-erythro-pentofuranosyl)-4-methoxy- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 154801-75-9 HCAPLUS
CN 1H-1,2,3-Triazolo[4,5-c]pyridine, 4-chloro-1-[5-[[[(1,1-dimethylethyl)diphenylsilyl]oxy]methyl]tetrahydro-2-furanyl]-, (2R-trans)-(9CI) (CA INDEX NAME)

RN 154801-76-0 HCAPLUS

CN 2-Furanmethanol, 5-(4-chloro-1H-1,2,3-triazolo[4,5-c]pyridin-1-yl)tetrahydro-, (2R-trans)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 154707-45-6

RL: BIOL (Biological study)

(preparation reaction with liquid ammonia)

RN 154707-45-6 HCAPLUS

CN 2H-1,2,3-Triazolo[4,5-c]pyridine, 4-chloro-2-(2-deoxy-β-D-erythropentofuranosyl)- (9CI) (CA INDEX NAME)

- L65 ANSWER 4 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN
- AN 1994:457887 HCAPLUS
- DN 121:57887
- ED Entered STN: 06 Aug 1994
- TI 2'-deoxy-2',2'-difluoro-(2,6,8-substituted) purine nucleosides having anti-viral and anti-cancer activity and intermediates
- IN Grindley, Gerald Burr; Grossman, Cora Sue; Hertel, Larry Wayne; Kroin,
 Julian Stanley
- PA Eli Lilly and Co., USA
- SO Eur. Pat. Appl., 20 pp.
 - CODEN: EPXXDW
- DT Patent
- LA English
- IC ICM C07H019-04

```
ICS A61K031-70
CC 33-9 (Carbohydrates)
```

Section cross-reference(s): 1, 63

FAN.CNT 1

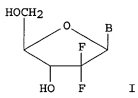
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE		
PI EP 576227	A2	19931229	EP 1993-304815	19930621 <		
EP 576227	A3	19940209				
R: AT, BE,	CH, DE, DK	, ES, FR,	GB, GR, IE, IT, LI,	LU, NL, PT, SE		
AU 9341347	A1	19931223	AU 1993-41347	19930618 <		
CA 2098876	AA /	19931223	CA 1993-2098876	19930621 <		
NO 9302287	Α	19931223	NO 1993-2287	19930621 <		
BR 9302433	Α	19940111	BR 1993-2433	19930621 <		
HU 64553	A2	19940128	HU 1993-1821	19930621 <		
JP 06056877	A2	19940301	JP 1993-149191	19930621 <		
CN 1084178	Α	19940323	CN 1993-107740	19930621 <		
PRAI US 1992-902304		19920622	<			
CLASS						
PATENT NO. CLA	SS PATENT	FAMILY CLA	ASSIFICATION CODES			

PATENT NO. CLASS PATENT FAMILY CLASSIFICATION CODES

EP 576227 ICM C07H019-04
ICS A61K031-70

OS MARPAT 121:57887

GI



- AB Title compds. I [B = purine, azapurine, deazapurine base] were prepared Thus, 2-amino-6-chloropurine was glycosidated, treated with MeNH2, and deblocked to give I [B = 2-amino-6-methylaminopurine] which had an IC50 against human leukemia cells of 0.054 μ g/mL and caused 56.9% inhibition of hepatitis B in vitro at 0.1 μ g/mL.
- ST deoxydifluororibofuranosylpurine nucleoside prepn antitumor virucide; purine deoxydifluororibofuranosyl
- IT Neoplasm inhibitors

Virucides and Virustats

(deoxydifluororibofuranosylpurine nucleosides)

IT Nucleosides, preparation

RL: SPN (Synthetic preparation); PREP (Preparation)

(deoxydifluororibofuranosylpurine, preparation and antitumor and virucidal activity of)

IT 156058-20-7P 156058-22-9P 156058-23-0P 156058-26-3P 156058-27-4P 156058-28-5P 156058-30-9P 156058-31-0P 156058-32-1P 156058-34-3P

156058-37-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and antitumor activity of)

IT 156058-29-6P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation and antitumor and virucidal activity of)

IT 155568-11-9P 155568-14-2P 156058-21-8P 156058-24-1P 156058-25-2P 156130-57-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent)

(preparation and reaction of, in preparation of

deoxydifluororibofuranosylpurine

nucleosides)

156058-33-2P 156058-35-4P 156058-36-5P TT 155568-11-9P

156058-41-2P 156058-38-7P 156058-39-8P 156058-40-1P

156058-46-7P 156058-43-4P 156058-44-5P 156058-45-6P 156058-42-3P 156124-75-3P

156058-48-9P 156100-12-8P 156124-74-2P 156058-47-8P

156124-76-4P **156124-77-5P** 156124-78-6P 156124-79-7P

156124-80-0P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

5451-40-1, 2,6-Dichloropurine 10310-21-1, 2-Amino-6-chloropurine TT

155131-42-3 153012-08-9

RL: RCT (Reactant); RACT (Reactant or reagent)

(reaction of, in preparation of deoxydifluororibofuranosylpurine nucleosides)

IT 156058-38-7P 156058-39-8P 156124-77-5P

156124-79-7P 156124-80-0P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 156058-38-7 HCAPLUS

1H-Imidazo[4,5-c]pyridine, 4,6-dichloro-1-(2-deoxy-2,2-difluoro-β-D-CNerythro-pentofuranosyl) - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

156058-39-8 HCAPLUS RN

1H-Imidazo[4,5-c] pyridin-4-amine, $1-(2-deoxy-2,2-difluoro-\beta-D-erythro-difluoro-\beta-D-erythro-difluoro-b-e$ CNpentofuranosyl) - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 156124-77-5 HCAPLUS

3H-Imidazo[4,5-b]pyridine, 5,7-dichloro-3-(3,5-di-0-benzoyl-2-deoxy-2,2-CN difluoro-β-D-erythro-pentofuranosyl) - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 156124-79-7 HCAPLUS

CN 1H-Imidazo[4,5-c]pyridine, 4-chloro-1-(3,5-di-O-benzoyl-2-deoxy-2,2-difluoro-β-D-erythro-pentofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 156124-80-0 HCAPLUS

CN 1H-Imidazo[4,5-c]pyridin-4-amine, 1-(3,5-di-0-benzoyl-2-deoxy-2,2-difluoro-β-D-erythro-pentofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L65 ANSWER 5 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1994:183004 HCAPLUS

DN 120:183004

```
ED
     Entered STN: 16 Apr 1994
     Therapeutic antiviral deoxythioribonucleosides
TI
     Koszalka, George Walter; Van Draanen, Nanine Agneta; Freeman, George
IN
     Andrew; Short, Steven Andersen; Slater, Martin John
     Wellcome Foundation Ltd., UK
PA
SO
     PCT Int. Appl., 39 pp.
     CODEN: PIXXD2
DT
     Patent
     English
LA
IC
     ICM A61K031-70
CC
     1-5 (Pharmacology)
     Section cross-reference(s): 33, 63
FAN.CNT 1
                        KIND
     PATENT NO.
                               DATE
                                           APPLICATION NO.
                                                                   DATE
                       ----
                                -----
                                            -----
                               19940120 WO 1993-GB1387
     WO 9401117
                          A1
PΙ
                                                                   19930701 <--
         W: AU, BB, BG, BR, CA, CZ, FI, HU, JP, KP, KR, KZ, LK, MG, MN, MW,
         NO, NZ, PL, RO, RU, SD, SK, UA, US, VN RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
             BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
     AU 9345084
                  A1 19940131 AU 1993-45084
                                                                   19930701 <--
PRAI GB 1992-14170
                                19920702 <--
     GB 1992-23181
                                19921105 <--
     WO 1993-GB1387
                                19930701 <--
CLASS
               CLASS PATENT FAMILY CLASSIFICATION CODES
 PATENT NO.
 ----- ,----
                        _____
 WO 9401117 ICM
                        A61K031-70
    MARPAT 120:183004
     2'-Deoxy-4'-thioribonucleosides and their physiol. acceptable salts,
     esters, or salts of such esters are useful for the manufacture of a medicament
    for the treatment or prophylaxis of retroviral, cytomegaloviral, varicella zoster viral, Epstein-Barr viral, human herpes virus 6, and hepatitis viral infections, including hepatitis B,
     coxsackie virus and hepatitis C virus infections.
     2'-Deoxy-4'-thioguanosine (preparation given) inhibited hepatitis B
     virus with an IC50 of <0.0032 \muM (74.5% inhibition) and a CCID50 of 13
     μM. Formulation examples are also given.
ST
     antiviral deoxythioribonucleoside; thiodeoxyribonucleoside virus
     inhibitor; deoxythioguanosine hepatitis B virus inhibitor
TT
     Virucides and Virustats
        (deoxythioribonucleosides)
IT
     Pharmaceutical dosage forms
        (of deoxythioribonucleosides, for treatment of virus infection)
TT
     Escherichia coli
        (trans-N-deoxyribosylase preparation from, for enzymic preparation of
antiviral
        deoxythioribonucleosides)
IT
     Virus, animal
        (Coxsackie, infection with, treatment of, with
        deoxythioribonucleosides)
IT
     Virus, animal
        (Epstein-Barr, infection with, treatment of, with
        deoxythioribonucleosides)
IT
    Virus, animal
        (cytomegalo-, infection with, treatment of, with
        deoxythioribonucleosides)
    Virus, animal
IT
        (hepatitis, infection with, treatment of, with
        deoxythioribonucleosides)
IT
    Virus, animal
        (hepatitis B, infection with, treatment of, with
```

deoxythioribonucleosides)

```
IT
     Virus, animal
        (hepatitis C, infection with, treatment of, with
        deoxythioribonucleosides)
IT
     Virus, animal
        (human cytomegalo-, inhibition of, with deoxythioadenosine)
IT
     Virus, animal
        (human herpes 6, infection with, treatment of, with
        deoxythioribonucleosides)
IT
     Pharmaceutical dosage forms
        (oral, of deoxythioribonucleosides, for treatment of virus infection)
TΤ
     Pharmaceutical dosage forms
        (parenterals, of deoxythioribonucleosides, for treatment of virus
        infection)
     Virus, animal
TT
        (retro-, infection with, treatment of, with deoxythioribonucleosides)
IT
     Virus, animal
        (varicella-zoster, infection with, treatment of, with
        deoxythioribonucleosides)
     9026-93-1, Adenosine deaminase
IT
     RL: BIOL (Biological study)
        (in preparation of antiviral deoxythioguanosine)
IT
     135656-41-6P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation and reaction of, in preparation of antiviral
deoxythioadenosine)
     153585-36-5P
IT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of)
ΙT
     135656-33-6P
                    153666-10-5P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of and virus infection inhibition with)
IT
     9026-86-2P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, from Escherichia coli, for enzymic preparation of antiviral
        deoxythioribonucleosides)
IT
     153585-34-3P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of, from aminomethoxypurine, trans-N-deoxyribosylase in)
     87-42-3, 6-Chloropurine
                               153585-35-4
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, in preparation of antiviral deoxythioadenosine)
IT
     134111-32-3
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with aminomethoxypurine, trans-N-deoxyribosylase in)
     20535-83-5, 2-Amino-6-methoxypurine
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reaction of, with deoxythiouridine, trans-N-deoxyribosylase in)
                                                               153585-22-9
IT
     153585-20-7
                   153585-20-7D, halo derivs.
                                                 153585-21-8
     153585-22-9D, halo derivs.
                                 153585-23-0
                                                 153585-24-1
                                                               153585-25-2
                   153585-26-3D, halo derivs.
                                                 153585-27-4
                                                               153585-28-5
     153585-26-3
     153585-28-5D, halo derivs.
                                 153585-29-6 153585-30-9
     153585-30-9D, halo derivs. 153585-31-0
     153585-32-1 153585-32-1D, halo derivs.
                                153666-08-1
                   153666-07-0
                                               153666-08-1D, halo
     153585-33-2
              153666-09-2 153666-10-5D, halo derivs.
     derivs.
    RL: BIOL (Biological study)
        (virus infection inhibition with)
     153585-30-9 153585-30-9D, halo derivs.
TT
     153585-31-0 153585-32-1 153585-32-1D, halo
     derivs. 153585-33-2
     RL: BIOL (Biological study)
```

(virus infection inhibition with)

RN 153585-30-9 HCAPLUS

CN 1H-Imidazo[4,5-c]pyridin-4-amine, 1-(2-deoxy-4-thio-D-erythro-pentofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 153585-30-9 HCAPLUS

CN 1H-Imidazo[4,5-c]pyridin-4-amine, 1-(2-deoxy-4-thio-D-erythro-pentofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 153585-31-0 HCAPLUS

CN 1H-Imidazo[4,5-c]pyridine-4,6-diamine, 1-(2-deoxy-4-thio-D-erythro-pentofuranosyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 153585-32-1 HCAPLUS

CN 4H-Imidazo[4,5-c]pyridin-4-one, 1-(2-deoxy-4-thio-D-erythro-pentofuranosyl)-1,5-dihydro- (9CI) (CA INDEX NAME)

RN 153585-32-1 HCAPLUS

CN 4H-Imidazo[4,5-c]pyridin-4-one, 1-(2-deoxy-4-thio-D-erythro-pentofuranosyl)-1,5-dihydro-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 153585-33-2 HCAPLUS

CN 4H-Imidazo[4,5-c]pyridin-4-one, 6-amino-1-(2-deoxy-4-thio-D-erythro-pentofuranosyl)-1,5-dihydro- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L65 ANSWER 6 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1992:584333 HCAPLUS

DN 117:184333

ED Entered STN: 15 Nov 1992

TI Nucleobase transporter-mediated permeation of 2',3'-dideoxyguanosine in human erythrocytes and human T-lymphoblastoid CCRF-CEM cells

AU Gati, Wendy P.; Paterson, Alan R. P.; Tyrrell, David L. J.; Cass, Carol E.; Moravek, Josef; Robins, Morris J.

CS Dep. Pharmacol., Univ. Alberta, Edmonton, AB, T6G 2H7, Can.

SO Journal of Biological Chemistry (1992), 267(31), 22272-6 CODEN: JBCHA3; ISSN: 0021-9258

```
DT
     Journal
LA
     English
CC
     1-5 (Pharmacology)
     Several 2',3'-dideoxynucleosides (ddNs), agents that inhibit the
AB
     replication of human immunodeficiency virus and hepatitis B
     virus, enter mammalian cells by simple diffusion. In this report, the
     authors show that the membrane permeation of 2',3'-dideoxyguanosine (ddG)
     in human erythrocytes and CCRF-CEM cells, in contrast with that of other
     ddNs, is transporter-mediated. Inward fluxes of ddG in both cell types
     were inhibited by adenine, hypoxanthine, and acyclovir, but not by
     inhibitors of nucleoside transport (nitrobenzylthioinosine, dipyridamole,
     dilazep). Fluxes of ddG in human erythrocytes were attributable to a
     single, rate-saturable process (Km, 380 \pm 90 \mu M and Vmax, 7.9 \pm
     0.8 pmol/s/\mu L cell water) that was competitively inhibited by adenine
     (Ki, 16 \mu M)\,. These results showed that ddG entered human erythrocytes
     and CCRF-CEM cells by a transporter-mediated process that was also the
     basis for entry of purine nucleobases. In contrast, inward fluxes of
     2,6-diaminopurine-2,3'-dideoxyriboside (ddDAPR), a prodrug of ddG, were
     not affected by purine nucleobases or nucleoside transport inhibitors in
     either cell type. Thus, the permeation properties of ddDAPR resembled
     those of 2',3'-dideoxyadenosine, a diffusional permeant (cell uptake is
     transporter-independent), and contrasted with those of ddG, the
     deamination product of ddDAPR. This study demonstrated that the
     nucleobase moiety of ddNs is an important determinant of membrane
     permeation.
ST
     dideoxyquanosine membrane permeation nucleobase transporter; erythrocyte
     nucleobase transport system dideoxyguanosine permeation; lymphoblast
     nucleobase transport system dideoxyguanosine permeation
IT
     Erythrocyte
        (dideoxyguanosine permeation in human, nucleobase transporter mediation
IT
     Virucides and Virustats
        (dideoxyguanosine, nucleobase transporter-mediated membrane permeation
        of, in human cells)
IT
    Partition
        (of dideoxynucleosides between octanol and buffer)
    Animal cell line
IT
        (CCRF-CEM, dideoxyguanosine permeation in human, nucleobase transporter
        mediation of)
    Biological transport
TT
        (permeation, of dideoxyguanosine in human erythrocytes and
        T-lymphoblastoid CCRF-CEM cells, nucleobase transporter mediation of)
     58-32-2, Dipyridamole 58-63-9, Inosine 68-94-0, Hypoxanthine
IT
     73-24-5, Adenine, biological studies 961-07-9, 2'-Deoxyguanosine
                              35898-87-4, Dilazep
                                                   38048-32-7
     13877-76-4, Formycin B
     59277-89-3, Acyclovir
     RL: BIOL (Biological study)
        (dideoxyquanosine transport by human erythrocytes and T-lymphoblastoid
        CCRF-CEM cells response to)
     85326-06-3, 2',3'-Dideoxyguanosine
IT
     RL: PROC (Process)
        (nucleobase transporter-mediated permeation of, in human erythrocytes
        and T-lymphoblastoid CCRF-CEM cells)
     4097-22-7, 2',3'-Dideoxyadenosine 69655-05-6, 2',3'-Dideoxyinosine
IT
     107550-73-2
     RL: BIOL (Biological study)
        (transport of, by human erythrocytes and T-lymphoblastoid CCRF-CEM
        cells, nucleobase transport system in relation to)
     13877-76-4, Formycin B
IT
     RL: BIOL (Biological study)
        (dideoxyguanosine transport by human erythrocytes and T-lymphoblastoid
        CCRF-CEM cells response to)
```

13877-76-4 HCAPLUS

RN

CN 7H-Pyrazolo[4,3-d]pyrimidin-7-one, 1,4-dihydro-3-β-D-ribofuranosyl-(8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry.

```
L65 ANSWER 7 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN
     1992:59866 HCAPLUS
AN
DN
     116:59866
     Entered STN: 21 Feb 1992
ED
     Synthesis and evaluation of anti-HIV-1 and antitumor activity of
TI
     2',3'-didehydro-2',3'-dideoxy-3-deazaadenosine, 2',3'-dideoxy-3-
     deazaadenosine and some 2',3'-dideoxy-3-deazaadenosine 5'-dialkyl
     phosphates
     Franchetti, P.; Cappellacci, L.; Cristalli, G.; Grifantini, M.; Pani, A.;
ΑU
     La Colla, P.; Nocentini, G.
     Dip. Sci. Chim., Univ. Camerino, Camerino, 62032, Italy
CS
     Nucleosides & Nucleotides (1991), 10(7), 1551-62
SO
     CODEN: NUNUD5; ISSN: 0732-8311
DT
     Journal
LA
     English
CC
     33-9 (Carbohydrates)
     Section cross-reference(s): 1
GI
```

The 4-amino-1-(2,3-dideoxy-β-D-glycero-pent-2-enofuranosyl)-1H-imidazo[4,5-c]pyridine (I), 4-amino-1-(2,3-dideoxy-β-D-glycero-pentofuranosyl)-1H-imidazo[4,5-c]pyridine (II), and 3-deaza analogs of the anti-HIV agents 2',3'-didehydro-2',3'-dideoxyadenosine and 2',3'-dideoxyadenosine, have been synthesized. The reaction of 3-deazaadenosine with 2-acetoxyisobutyryl bromide yielded a mixture of cis and trans 2',3'-halo acetates which was converted into olefinic nucleoside I on treatment with a Zn/Cu couple and then with NH3/MeOH. A number of

phosphate triester derivs. of II have also been prepared Nucleotides III (R = Et, Pr, Bu) and 3-deazaadenosine have shown anti-HIV activity at non-cytotoxic doses. III have also shown significant cytostatic activity against murine colon adenocarcinoma cells. deoxydeazaadenosine phosphate virucide neoplasm inhibitor; dehydrodideoxyadenosine prepn virucide neoplasm inhibitor; deazaadenosine didehydrodideoxy virucide neoplasm inhibitor; nucleotide dideoxydeaza virucide neoplasm inhibitor Neoplasm inhibitors Virucides and Virustats (dideoxydeazaadenosine phosphates as)

ΙT Nucleotides, polymers

ST

IT

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(di-, deoxy-, deazaadenine, preparation and antiviral and antitumor activity

Nucleotides, biological studies IT

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(dideoxy-, deazaadenine, preparation and antiviral and antitumor activity of)

4097-22-7 7057-48-9 TТ

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(antiviral and antitumor activity of)

819-43-2 2510-89-6 TT 814-49-3

RL: RCT (Reactant); RACT (Reactant or reagent) (phosphorylation by, of dideoxydeazaadenosine)

138352-56-4P 138352-57-5P 138352-58-6P IT

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and antiviral and antitumor activity of)

138352-55-3P IT

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation, hydrogenation, antiviral and antitumor activity of)

IT 130948-34-4P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation, phosphorylation, antiviral and antitumor activity of)

ΙT 40635-67-4

RL: RCT (Reactant); RACT (Reactant or reagent)

(reaction of, with deazaadenosine)

ΙT 6736-58-9

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(sequential bromination and elimination reaction, antiviral, and antitumor activity of)

138352-56-4P 138352-57-5P 138352-58-6P ΙT

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and antiviral and antitumor activity of)

138352-56-4 HCAPLUS RN

Phosphoric acid, [5-(4-amino-1H-imidazo[4,5-c]pyridin-1-yl)tetrahydro-2-CN furanyl]methyl diethyl ester, (2S-cis) - (9CI) (CA INDEX NAME)

RN 138352-57-5 HCAPLUS

CN Phosphoric acid, [5-(4-amino-1H-imidazo[4,5-c]pyridin-1-yl)tetrahydro-2-furanyl]methyl dipropyl ester, (2S-cis)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 138352-58-6 HCAPLUS

CN Phosphoric acid, [5-(4-amino-lH-imidazo[4,5-c]pyridin-1-yl)tetrahydro-2-furanyl]methyl dibutyl ester (9CI) (CA INDEX NAME)

IT 138352-55-3P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation, hydrogenation, antiviral and antitumor activity of)

RN 138352-55-3 HCAPLUS

CN 2-Furanmethanol, 5-(4-amino-1H-imidazo[4,5-c]pyridin-1-yl)-2,5-dihydro-, (2S-cis)- (9CI) (CA INDEX NAME)

IT 130948-34-4P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation, phosphorylation, antiviral and antitumor activity of)

RN 130948-34-4 HCAPLUS

CN 2-Furanmethanol, 5-(4-amino-1H-imidazo[4,5-c]pyridin-1-yl)tetrahydro-, (2S,5R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 6736-58-9

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(sequential bromination and elimination reaction, antiviral, and antitumor activity of)

RN 6736-58-9 HCAPLUS

CN 1H-Imidazo[4,5-c]pyridin-4-amine, 1- β -D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L65 ANSWER 8 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1990:544908 HCAPLUS

DN 113:144908

ED Entered STN: 27 Oct 1990

- ΤI Inhibition of hepatitis A virus replication in vitro by antiviral compounds Crance, J. M.; Biziagos, E.; Passagot, J.; Van Cuyck-Gandre, H.; Deloince, ΑU
- Unite Biol. Mol., Cent. Rech. Serv. Sante Armees, La Tronche, 38702, Fr. CS Journal of Medical Virology (1990), 31(2), 155-60 SO

CODEN: JMVIDB; ISSN: 0146-6615

- DTJournal
- LΑ English
- CC 1-5 (Pharmacology)
- Forty antiviral compds. were screened for inhibitory effect on AB hepatitis A virus (HAV) antigen expression in the human hepatoma cell line PLC/PRF/5. Ribavirin, amantadine, glycyrrhizin, and pyrazofurin were selected in this screening test and were studied further. The selectivity indexes of these four compds., calculated as the ratio of 50% cytotoxic dose (determined by the trypan blue exclusion and by inhibition of [3H] leucine incorporation) to the 50% ED (determined by the viral antigen expression), were 4.6 and 3.0 with ribavirin, 5.3 and 5.9 with amantadine, 15.2 and 16.9 with glycyrrhizin, and 45.4 and 74.6 with pyrazofurin. All four compds. resulted in concentration-dependent redns. of

HAV antigen expression and HAV infectivity. Ribavirin, amantadine, pyrazofurin, and glycyrrhizin emerged, from the present study, as promising candidates for chemotherapy of acute hepatitis A.

- antiviral hepatitis A virus; ribavirin antiviral ST hepatitis A virus; amantadine antiviral hepatitis A virus; glycyrrhizin antiviral hepatitis A virus; pyrazofurin antiviral hepatitis A virus
- IT Virucides and Virustats

(against hepatitis A virus, screening for, in human hepatoma cells)

ITSaponins

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES

(antiviral activity of, against hepatitis A virus, in human hepatoma cells)

IT Virus, animal

> (hepatitis A, infection with, antiviral screening for therapy of, in human hepatoma cells)

TТ Pentosans

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES

(sulfates, antiviral activity of, against hepatitis A virus, in human hepatoma cells)

50-02-2, Dexamethasone 50-23-7, Hydrocortisone 50-81-7, Ascorbic acid, 54-25-1, 6-Azauridine biological studies 54-21-7, Sodium salicylate 58-08-2, Caffeine, biological studies 58-32-2, Dipyridamole 66-81-9, 73-03-0, Cordycepin 85-31-4, 6-Mercaptoguanosine Cycloheximide 113-00-8, Guanidine 141-84-4, Rhodanine 154-23-4, Catechin 378-44-9, Betamethasone 480-18-2, Taxifolin 320-67-2, 5-Azacytidine 1024-99-3, 5-Iodouridine 1123-54-2, 8-Azaadenine 768-94-5, Amantadine 1397-89-3, Amphotericin B 1405-86-3, 1147-23-5, 5-Iodocytidine 6990-06-3, Fusidic acid Glycyrrhizin 1445-07-4, Pseudouridine 9005-49-6, Heparin, biological studies 9042-14-2, 6998-60-3, Rifamycin 9072-19-9, Fucoidan 11089-65-9, Tunicamycin Dextran sulfate 13292-46-1, Rifampicin 13877-76-4 23205-42-7, 3-Deazauridine 26001-38-7, 8-Mercaptoguanosine 30868-30-5, Pyrazofurin 36791-04-5, Ribavirin RL: BAC (Biological activity or effector, except adverse); BSU (Biological

study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(antiviral activity of, against **hepatitis** A virus, in human hepatoma cells)

IT 13877-76-4 36791-04-5, Ribavirin

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(antiviral activity of, against **hepatitis** A virus, in human hepatoma cells)

RN 13877-76-4 HCAPLUS

CN 7H-Pyrazolo[4,3-d]pyrimidin-7-one, 1,4-dihydro-3-β-D-ribofuranosyl-(8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 36791-04-5 HCAPLUS

CN 1H-1,2,4-Triazole-3-carboxamide, 1- β -D-ribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L65 ANSWER 9 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1984:603955 HCAPLUS

DN 101:203955

TI Broad-spectrum synergistic antiviral activity of selenazofurin and ribavirin

AU Kirsi, Jorma J.; McKernan, Patricia A.; Burns, Noah J., III; North, James A.; Murray, Byron K.; Robins, Roland K.

CS Dep. Microbiol., Brigham Young Univ., Provo, UT, 84602, USA

SO Antimicrobial Agents and Chemotherapy (1984), 26(4), 466-75 CODEN: AMACCQ; ISSN: 0066-4804

DT Journal

LA English

CC 1-5 (Pharmacology)

The antiviral effects of selenazofurin [83705-13-9], ribavirin [36791-04-5], and 3-deazaguanosine [56039-11-3] were investigated sep. and in various combinations in an in vitro study. The combination interactions were evaluated at seven drug concns., graphically (isobolograms) or by using fractional inhibitory concentration indexes against mumps, measles, parainfluenza virus type 3, vaccinia and herpes simplex virus type 2 viruses in Vero and HeLa cells.

Selenazofurin in combination with ribavirin produced the greatest synergistic antiviral activity. However, the degree of synergy depended on the virus and cell line used. contrast, selenazofurin combined with 3-deazaguanosine consistently yielded an indifferent or an antagonistic response, or both, whereas the ribavirin-3-deazaguanosine interaction was additive against the same viruses. Single-drug cytotoxicity was minimal for the cytostatic agents selenazofurin and ribavirin but was markedly higher for cytocidal 3-deazaguanosine, as determined by relative plating efficiency after drug exposure. The drug combinations did not significantly increase cytotoxicity (they were only additive) when used on uninfected cells. Therefore, the enhanced antiviral activities of the drug combinations (shown to be synergistic) were due to specific effects against viral replication. These results indicated that in Vero and HeLa cells (i) the combination of selenazofurin and ribavirin produced and enhanced antiviral effect, thus requiring smaller amts. of drug to cause the same antiviral effect relative to a single compound; (ii) selenazofurin when compared with ribavirin and -deazaguanosine appeared to have a somewhat different mode of antiviral action; (iii) 3-deazaguanosine combined with selenazofurin was an unsuitable antiviral combination; and (i.v.) the antiviral activity of 3-deazaguanosine appeared to be due largely to its general overall cytotoxic effect. antiviral ribavirin selenazofurin combination;

deazaguanine antiviral combination

IT 36791-04-5

ST

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(antiviral activity of deazaguanine or selenazofurin and,

broad-spectrum synergistic)

IT 83705-13-9

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(antiviral activity of ribavirin or deazaguanine and,

broad-spectrum synergistic)

IT 56039-11-3

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(antiviral activity of ribavirin or selenazofurin and,

broad-spectrum synergistic)

IT 36791-04-5

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(antiviral activity of deazaguanine or selenazofurin and,

broad-spectrum synergistic)

RN 36791-04-5 HCAPLUS

CN 1H-1,2,4-Triazole-3-carboxamide, 1-β-D-ribofuranosyl- (9CI) (CF
INDEX NAME)

Absolute stereochemistry.

IT 56039-11-3

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(antiviral activity of ribavirin or selenazofurin and, broad-spectrum synergistic)

RN 56039-11-3 HCAPLUS

CN 4H-Imidazo[4,5-c]pyridin-4-one, 6-amino-1,5-dihydro-1-β-Dribofuranosyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

```
L65 ANSWER 10 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN
AN
     1969:46065 HCAPLUS
DM
     70:46065
ED
     Entered STN: 12 May 1984
TI
     Chemotherapeutic studies on mouse hepatitis virus. III.
     Antiviral effect of some pharmacodynamic drugs
AU
     Kanoh, Seizaburo
     Nat. Inst. Hyg. Sci., Osaka, Japan
CS
SO
     Chemotherapy (Tokyo) (1968), 16(6), 789-91
     CODEN: NKRZAZ; ISSN: 0009-3165
DT
     Journal
LΑ
     English
CC
     15 (Pharmacodynamics)
AB
     Xenaldial (100 μg./ml.) completely inactivated mouse hepatitis
     virus EHF-120 in vitro, but this drug (125 mg./kg./day, s.c., on the 1st 3
     days of infection) did not have an antiviral effect in mice. Treatment
     with s.c. injections of formicin (10-20 mg./kg./day) or
     p-carboxy-N-methylacetylnicotinic acid (?) (125-500 mg./kg./day) on the
     1st 3 days of infection decreased the mortality of infected mice from a
     control level of 57 to 42.8 and 37.5%, resp., in 1 group of mice.
     Treatment of mice with s.c. injections of Benadryl-HCl (0.5 mg./kg./day,
     for 4 days, beginning 2 days before infection) decreased the mortality
     rate from a control level of 43 to 28.5%.
ST
     antiviral effect formicin; formicin antiviral effect; xenaldial
     hepatitis virus; hepatitis virus nicotinates; virus
     hepatitis nicotinates; nicotinates hepatitis virus;
     Benadryl hepatitis virus control
```

IT Virucides

IT Viruses, animal

(mouse hepatitis, infection with, pharmaceuticals effect on)

IT 59-67-6, Nicotinic acid 147-24-0 1094-85-5 6742-12-7
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(virucidal activity of)

IT 6742-12-7

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(virucidal activity of)

RN 6742-12-7 HCAPLUS

CN D-Ribitol, 1-C-(7-amino-1H-pyrazolo[4,3-d]pyrimidin-3-yl)-1,4-anhydro-, (1S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

=>